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(71)Applicant: MARINE BIOTECHNOL INST CO

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(72)Inventor: IDENO AKIRA

MARUYAMA TADASHI

(54) METHOD FOR SUPPRESSING DENATURATION OF PROTEIN

(57) Abstract:

PROBLEM TO BE SOLVED: To provide a method for suppressing the denaturation of a protein by heat or an organic solvent to prevent the irreversible formation of aggregates. SOLUTION: This method for suppressing the denaturation of a protein comprises making FKBP type PPIase that is derived from a thermophilic or hyperthermophilic archaea and has a molecular weight from 26 to 33 kDa coexist with the protein.

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CLAIMS

[Claim(s)]

[Claim 1] In the amino acid sequence of the protein (b) array number 2 expressed according to the amino acid sequence of the protein (a) array number 2 shown in the following (a) – (c), or four publications, or four publications 1 or two or more amino acid are expressed by deletion and the amino acid sequence permuted or added. According to and the base sequence of the protein (c) array number 1 which has PPIase activity or chaperon Mr. activity, or three publications Protein which DNA with which it is expressed or it and complementary DNA, and DNA hybridized under stringent conditions are protein of the super—thermophilic nature or the thermophile Archea origin which carries out a code, and has PPIase activity or chaperon Mr. activity [claim 2] protein according to claim 1 is coexisted with other protein — making — being concerned — others — the denaturation restrainning of the protein characterized by controlling proteinic denaturation.

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DETAILED DESCRIPTION

Detailed Description of the Invention

Field of the Invention] This invention relates to the approach of controlling the denaturation of the protein produced by environmental stress, such as heat. Description of the Prior Art.] Protein is the polypeptide with which two or more amino acid was protein production by gene modification technology, purpose protein is produced as an inclusion tertiary structure (spacial configuration) formed of the interaction between intramolecular or a body with the inactive protein overproduced in host bacteria, such as Escherichia coli, and the environmental variation, it is always mentioned as a technical problem. Moreover, in the useful connected by peptide linkage. In order for protein to discover the property, the characteristic molecule is important. Generally, if protein adds environmental stressors, such as heat, the spacial configuration will change and the property will disappear irreversibly in many cases. Therefore, when how protein is maintained at a stable condition treats protein to such an technical problem that productive efficiency falls is mentioned.

has the characteristic structure that the doughnut mold structure where seven subunits stood in change, it is produced. These exist widely regardless of a procaryote and eukaryote, and GroE is solution contain nucleotides, such as SHAPERONIN protein and ATP, and stabilizing the enzyme SHAPERONIN has and a spacial configuration deforms them with heat or a modifier, they attains a row annularly lapped with two steps and of consisting of a total of 14 subunits. GroEL catches [0004] When each of these uses a spacial configuration formation operation of the protein which the purpose according to the operation which rewinds a proteinic polypeptide chain to an original in a solution" is proposed by JP,7-67641,A. Moreover, "the approach using SHAPERONIN which which denaturalized chemically, genetic manipulation, etc. from Thermus thermophilus (Thermus does not choose a proteinic class but it is shown clearly that it participates in proteinic spacial configuration formation nonspecific. For example, GroEL which is the construct of Above GroE [0003] Research is energetically made as a solution of the above-mentioned trouble until now, and various improvement proposals are proposed. In recent years, the interest is increasing in denatured protein to the crevice of doughnut structure, and folding up to the protein of a right formation and a proteinic structural change. A molecular chaperone is a group of a heat shock well known as a molecular chaperone produced [especially] from Escherichia coli. This GroE spacial configuration efficiently is known with consumption of nucleotides, such as ATP, and thermophilus) for the purpose of reproducing activity protein" is proposed by JP,7-48398,A. protein, and when a cell is exposed to various environmental stress, such as a temperature refined the inactive protein accumulated into the transformant used by the inactive protein proteinic stabilization are made. For example, "the approach of making an enzyme content association of GroES which is cofactor. Some attempts which apply this SHAPERONIN to the molecular chaperone as a factor which participates in proteinic spacial configuration

live together generally. in order that [moreover,] SHAPERONIN may act by purpose protein and [0005] However, SHAPERONIN needs to make high energy matter, such as ATP, CPT, and UDP, spacial configuration (it folds up).

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the mole fraction of 1:1 -- molecular weight -- 1 million -- it is necessary to use very highconcentration SHAPERONIN, and economical efficiency is missing in SHAPERONIN of the amount of macromolecules near Da(s).

attention to the FKBP type PPlase of the Archea origin. Consequently, it has been shown clearly trans isomerization reaction of the amino terminal side peptide linkage of the proline residue in a each immunosuppresant. Moreover, the Per Boleyn (Parvulin) type with which the homologies of polypeptide chain, it was expected as what can be used for proteinic stabilization or playback of the target molecule of FK506 which are known as an immunosuppresant, and is divided roughly into a cyclo FIRIN type and a FKBP (FK506 binding protein) type from the susceptibility over and the activity (chaperon Mr. activity) which controls proteinic irreversible condensation in an FrontBiosoi.2000 Sep 1, 5, D 821-836). Unlike the above-mentioned chaperon, PPlase has the 0006] On the other hand, PPlase (Petidyl prolyl cis-trans isomerase) is the cyclosporine and that it has not only the above-mentioned PPlase activity but the activity which increases the yield of the protein volume return originally considered as the function of SHAPERONIN itself an amino acid sequence also differ by the insusceptibility in both immunosuppresant is being reproduction speed of protein higher order structure by carrying out the catalyst of the cisinactive protein like above-mentioned SHAPERONIN. Artificers have so far inquired paying found in recent years. Since PPlase had the function (PPlase activity) to promote the interesting thing at the FKBP type PPlase of these origins (Maruyama et al., 2000, advantage that high energy matter called ATP etc. is unnecessary.

[0007] Furthermore, although, as for PPlase marketed until now, Kamiichi of two kinds of a cyclo of the animal origin is very weak with heat. On the other hand, PPlase of the Archea origin does and needed to be saved at the low temperature of 4 degrees C or -20 degrees C. Thus, PPIase sigma company, respectively, all were the things of the animal origin, and the stability was low not lose activity, even if it is strong with heat and long duration neglect is carried out under a high temperature service. For this reason, that use is expected as a PPlase reagent with new FIRIN type and FKBP type things [every one kind of] was carried out from for example, the PPlase of the Archea origin.

terminal domain part, and the remaining C terminal domain part is PPlase of 2 functionality which domain of kDa type PPlase is the high field of a thing short type and homology, compared with a Problem(s) to be Solved by the Invention The great portion of PPIase of the Archea origin is a (Maruyama, T and Furutani, M Front Biosci. 2000 Sep 1, 5, D821-836; Jida et al., 2000, and Gene weight is 17-18 kDa extent, and the thing long type type molecular weight is 26-33 kDa extent (Methanococcus thermoautotrophicum) origin PPlase activity is weak, for example, is [origin] FKBP type. Moreover, FKBP type PPlase is classified into the thing short type type molecular 256,319-326), the FKBP type PPlase which consists of 26-33 kDa of the Archea origin -- the such a technological background, and aims at offering 25-35 kDa type highly efficient PPIase. shows high efficiency is found out, the utility value is very high. This invention is made under (Ideno et al., 2000, Eur.J.Biochem 267, 3139-3148). 25-35 Also in kDa type PPlase, if what thermophile Archea shows only the activity of or less 1/1000 extent of 16kDa type PPlase. about 17 to 18 kDa from -- the field which bears PPIase activity is in the becoming amino has the function which controls proteinic condensation. 26–33 Although the amino terminal 17-18kDa type thing, PPIase of the methano KOKKASU thermostat auto trophy cam

Means for Solving the Problem] this invention person completed a header and this invention for origin belongs to a 26-33 kDa type as a result of repeating examination wholeheartedly, in order (Pyrococcus horikoshii) and the METANOKOKKASU YANASHII (Methanococcus jannaschiii) having very strong chaperon Mr. activity, although PPlase of pie ROKOKKASU HORIKOSHI to solve the above-mentioned technical problem.

[0010] That is, this invention is protein shown in the following (a) – (c).

amino acid sequence the array number 2 or given in four, or four publications 1 or two or more (a) In the amino acid sequence of the protein (b) array number 2 expressed according to an amino acid are expressed by deletion and the amino acid sequence permuted or added.

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According to and the base sequence of the protein (c) array number 1 which has PPlase activity PPlase activity or chaperon Mr. activity, and this invention coexist the above-mentioned protein with other protein -- making -- being concerned -- others -- it is the denaturation restrainning the thermophile Archea origin in which DNA with which it is expressed or it and complementary or chaperon Mr. activity, or three publications It is protein of the super-thermophilic nature or DNA, and DNA hybridized under stringent conditions carry out a code, the protein which has of the protein characterized by controlling proteinic denaturation.

Embodiment of the Invention] Hereafter, this invention is explained to a detail. The protein of this invention contains the protein shown in the following (a) – (c).

According to and the base sequence of the protein (c) array number 1 which has PPlase activity amino acid sequence the array number 2 or given in four, or four publications 1 or two or more (a) In the amino acid sequence of the protein (b) array number 2 expressed according to an amino acid are expressed by deletion and the amino acid sequence permuted or added.

DNA, and DNA hybridized under stringent conditions carry out a code. Each protein of protein (a) (c) which has PPlase activity or chaperon Mr. activity is FKBP type PPlase of 26-33kDa. The the thermophile Archea origin in which DNA with which it is expressed or it and complementary protein of (a) is PPlase of pie ROKOKKASU HORIKOSHI or the METANOKOKKASU YANASHII or chaperon Mr. activity, or three publications It is protein of the super-thermophilic nature or

variation includes an artificial variation besides the variation produced in a nature. As a means to 43, 1101-1111). On the other hand, chaperon Mr. activity can use rhodanese, citrate synthetase, preferably, is less than ten amino acid still more preferably, and is less than five amino acid most a malate dehydrogenase, glucose-6-phosphate dehydrogenase, etc. as a model enzyme (Kawada produce artificial variation, although a site-directed-mutagenesis method (Nucleic Acids Res.10, 6487-6500, 1982) etc. can be mentioned, it is not necessarily limited to this. Unless the number preferably. Chymotrypsin couple DOASSEI which Fischer and others proposed can estimate the 1998, bioscience, an industry 56, and 593-598), and can evaluate them by the regeneration rate PPlase activity as used in the field of this invention (Fischer et al., 1984, Biomed.Biochim.Acta PPlase activity or chaperon Mr. activity lose was introduced into the protein of (a). Such a [0012] The protein of (b) is protein with which the variation of extent which does not make of the varied amino acid makes PPlase activity or chaperon Mr. activity lose, although the number is not restricted, usually it is less than 30 amino acid, is less than 20 amino acid

guanidine hydrochloride. As an approach Jolo Bitsch's and others approach evaluates (Horowitz, and others approach (Taguchi et al.1994, J.Biol.Chem.269, 8529-8534) etc. is mentioned to the 1995, Methods Mol.Biol.40,361-368), and condensation control of denatured protein, Taguchi's of denatured protein and the rate of control of condensation of denatured protein which are SHAPERONIN for these after denaturation processing with protein modifiers, such as 6M started when a modifier is diluted with the buffer solution which contains PPlase and approach of evaluating the regeneration rate of denatured protein, respectively.

[0013] The protein of (c) is PPiase of thermophile Archea obtained by using the hybridization of means conditions in which only specific hybridization occurs and nonspecific hybridization does not occur. Usually such conditions are "IxSSC, 0.1%SDS, 37-degree-C" extent, are "0.5xSSC, 0.1%SDS, 42-degree-C" extent preferably, and are "0.2xSSC, 0.1%SDS, 65-degree-C" extent DNA, or the super-thermophile Archea origin. "The stringent conditions" in the protein of (c) according to the base sequence of array number 1 or array number 3 publication, and usually high homology, high homology --- 60% or more of homology --- desirable --- 75% or more of still more preferably. DNA obtained by hybridization has DNA with which it is expressed homology -- 90% or more of homology is pointed out still more preferably.

thermophile Archea or super-thermophile Archea is cultivated, and genomic DNA is extracted for [0014] The protein of this invention is obtained by the following approaches, for example. First, bacteriolysis using bacillus solubilizing agents, such as SDS. As the super-thermophilic nature the obtained fungus body by technique, such as a phenol extract and ethanol settling, after a used by this invention, or thermophile Archea An ASHIDI anus (Acidianus) group, a meta-loss

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group and a cardo KOKKASU (Caldococcus) group, AKIOGUROBA loess (Archaeoglobales) eyes, group, A methano sir mass (Methanothermus) group, the Methanobacterium (Methanobacterium) KOKKARESU (Thermococcales) eyes, such as a methano piler loess (Methanopyrales) eye, a pie IGUNEOKOKKARESU (Igneococcules) eyes, such as a hyper-Thermus (Hyperthermus) group, a methano cardo KOKKASU (Methanocaldococcus) group and a meta-noy GUNISU (Methanoignis) (Methanosarcinales) eyes, such as a methano ZARUCHINA (Methanosarcina) group, Thermostat ROKOKKASU (Pyrococcus) group, and a thermostat KOKKASU (Thermococcus) group, Archea, Thermostat Protea loess (Thermoproteales) eyes, such as a thermostat FIRAMU (Thermofilum) use the thing of the super-thermophile Archea origin also in these Archea, and it is desirable to (Thermoplasma) group and a PIKURO philus (Picrophilus) group, is mentioned. It is desirable to BAKUTA (Methanothermobacter) group and a methano SUFAERA (Methanosphaera) group, A (Sulfiolobus) group, Sulfo ass loess (Sulfolobales) eyes, such as a SURUFUROKOKKASU (Sulfurococcus) group and a SURUFURISUFAERA (Sulfurisphaera) group, An AERO pie ram Stetteria) group, a SUTAFIRO Thermus (Staphylothermus) group, A thermostat DISUKASU such as an AKIOGUROBUSU (Archaeoglobus) group and a ferro GUROBUSU (Ferroglobus) (Methanothermococcus) group, Methano KOKKARESU (Methanococcales) eyes, such as a group, A methano micro BAIARESU (Methanomicrobiales) eye, Methano ZARUCHINARESU (Aeropyrum) group, a DESURUFUROKOKKASU (Desulfurococcus) group, A SUTETTERIA FAERA (Metallosphaera) group, A stage OROBASU (Stygiolobus) group, a sulfo ROBASU group, Methano bacteria loess (Methanobacteriales) eyes, such as a methano thermostat pie RODIKUTIUMU (Pyrodictium) group, and a pie ROROBASU (Pyrolobus) group, A pie ROBAKYURAMU (Pyrobaculum) group, a thermostat Proteus (Thermoproteus) group, (Thermodiscus) group, an IGUNEOKOKKASU (Igneococcus) group, A Sir MOS FAERA such as thermoplasma loess (Thermoplasmales) eyes, such as a thermostat plasma (Thermosphaera) group and a sulfo FOBOKOKKASU (Sulfophobococcus) group, methano KOKKASU (Methanococcus) group, a methano thermostat KOKKASU

phagemid DNA, such as lambdaZAP, the various vectors 10, for example, lambdagt, of the lambda [0015] Thus, the obtained genomic DNA is connected with a suitable vector after cutting with a [0016] On the other hand, DNA which is equivalent to it based on the amino acid sequence of suitable restriction enzyme, and a genomic DNA library is produced. The plasmid vector of phage origin, or a pUC18 and pBR322 grade can be used for a vector.

especially use Archea of a pie ROKOKKASU group and Methanococcus.

fragment of the gene which carries out the code of the protein of this invention can be obtained. different living thing kind origin, and it considers as the primer used for PCR. As an example of The above-mentioned partial fragment can be used as a probe of a gene screening by carrying amino acid sequence of 137 to 144 number, etc. can be mentioned, for example. If PCR which uses the above-mentioned genomic DNA as mold using such a primer is performed, the partial the high field of a homology, the 25-32nd amino acid sequences in PPlase of drawing 1, the out an indicator with radioelements, such as [32P], and nonradioactive compounds, such as the high field of a homology is compounded between the FKBP type PPlase genes of the JIKOKISHI genin.

gene which carries out the code of the protein of this invention. It can opt for the decision of a With the above procedure, all the DNA sequences that carry out the code of the protein of this invention containing a termination codon can be isolated from a translation initiation codon. It is [0017] What is necessary is to introduce the above-mentioned genomic DNA library into hosts, possible to carry out extensive preparation of the protein of this invention by inserting suitably DNA which carries out the code of the isolated protein of this invention in expression vectors, base sequence by the Sanger's method or the general approach of the maxim-gilbert method. mentioned probe which carried out labeling, in order to acquire all the base sequences of the such as a pET system, introducing into a microorganism or a cultured cell and making them such as Escherichia coli, and just to choose the clone strongly combined with the above discovered by the above-mentioned actuation.

0018] What is necessary is just to make target protein coexist with the protein of this invention or denaturation control of unstable protein. It not only excels in thermal resistance, but the

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protein of this invention demonstrates depressor effect to proteinic various denaturation in order to show the outstanding cold resistance, organic solvent-proof nature, etc. As a means made to ratio of 0.1-500 is suitable for it to purpose protein 1. Moreover, coexpression should just insert Conversely, even if it builds purpose protein into the pACYC system and builds the PPlase gene invention is excellent in thermal resistance, and advantageous in the cases, such as conveyance this invention with DNA which carries out the code of the target protein, etc. can be mentioned to target protein, for example. When mixing with target protein, the ratio is a mole ratio and the together within a host. The manifestation of both gene is attained under each promotor control. inclusion body. If the protein of this invention is coexpressed, while the denaturation of purpose in the downstream of the manifestation promoter of for example, a pACYC system plasmid the live together, a means to mix the protein of this invention, a means to coexpress the protein of activity and chaperon Mr. activity, and it is produced as a soluble fraction. The protein of this protein will be controlled, inclusion body formation is barred by effectiveness, such as PPlase in CoIE1 system, it does not interfere. If this coexpression system is used, when discovering gene which carries out the code of the protein of this invention. For example, when purpose protein is inserted in the expression vector with the DNA replication initiation field of coIE1 bacterium, if another drug resistance marker is made to own, both gene lives each plasmid solubilization fraction produce the protein discovered as denaturation objects, such as an systems, such as pET, since a pACYC vector can live together within them and a host purpose protein with Escherichia coli etc., for example, it becomes possible to make a

and preservation. [0019]

[Example] Hereafter, although an example explains this invention, the range of this invention is not limited to this.

[Example 1] Cloning of the preparation PPlase gene of the genomic DNA of super-thermophile Archea was carried out from pie ROKOKKASU HORIKOSHI and METANOKOKKASU YANASHII. Pie ROKOKKASU HORIKOSHI (MD151) came to hand from the Institute of Physical and Chemical Research, and received METANOKOKKASU YANASHII from DSM.
[0020] It is the bacillus suspension of pie ROKOKKASU YANASHII from DSM.
[0020] It is the bacillus suspension of pie ROKOKKASU HORIKOSHI and METANOKOKKASU YANASHII, respectively About 50 mul extraction was carried out and fungus bodies were collected by carrying out at-long-intervals alignment separation for 15000 pmx 5 minutes. After 500micro of TE buffer solutions I washed the fungus body twice, it suspended in TE buffer solution which contains SDS 0.1%, and bacteriolysis processing was performed for 60 minutes at 95 degrees C. After equivalent TE buffer-solution saturation phenol and chloroform performed deproteinization processing, each genomic DNA was settled in ethanol.

Deproteinization processing, each genomic DNA was settled in ethanol.

[0021] [Example 2] Based on the array information on the pie ROKOKKASU HORIKOSHI origin PPlases shown in magnification, cloning drawing 1 and drawing 2 of a PPlase gene, and the array information on the METANDKOKKASU YANASHII origin PPlase, the PPlase (it is hereafter written as "PHFK") gene of the pie ROKOKKASU HORIKOSHI origin and the PPlase (it is bereafter written as "MJFKL") gene of the METANDKOKKASU YANASHII origin were amplified by the PCR method, As a primer for gene amplification, about PHFK, PHFK-F1 and PHFK-R1 were used about MJLFK, respectively (Table 1). The restriction enzyme site was established in these primers. Moreover, a reaction presentation and reaction cycle of PCR are as being shown in the table 2 *** table 3, respectively. DNA

polymerase used TaKaRaEx.Taq.

Table 1

アンダーライン: 各制段酵素サイト

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60/60/40

[UUZ3] [Table 2] PCRの反応組成

I ON DODAHAMA	
Reaction buffer($\times 10$)	1041
GLATE	8 44 1
Ex Taq	0.5 41
ナノム DNA(10ng/μ1)	2 m l
Forward Primer(20pmol/ μ l)	4μ1
Reverce Primer(20pmol/ μ 1)	4μl
滅酷水	71.5 41
4年	1007

[0024]

[Table 3] PCRの反応条件

1 Icycle	nin	30cycle	
95°C×5min	96°C×0. 5min	59°C×1min	72°C × Imin
ブレヒート	変性	アニーリング	海南

[0025] The band part which includes after separation each magnification product acquired by the PCR method for a DNA fragment by agarose gel electrophoresis 2% was started, and phenol chloroform processing and ethanol precipitate extracted the purpose DNA. A DNA fragment is dissolved in sterilized water and it is the pT7 blue T plasmid vector of the amount of 10 times to each about ten to 100 ng. (Novegen) in addition, ligation of the DNA fragment was carried out by processing at 16 more degrees C for 1 hour.

flask of 2L. (Yeast Extract 16 g L-1, BACTO TRYPTON 20 g L-1, NaCl5g L-1, ampicillin 100 mug the primer corresponding to a DNA fragment was made into the electropositive colony overnight. Ligation was carried out, pET21a plasmid DNA is collected from the electropositive colony which contains a PPlase gene after carrying out transformation by adding the obtained ligation reaction acquired PCR product was determined by using a plasmid as mold and performing the sequence [0028] [example 4] The manifestation of PHFK and MJFKL was tried using the purification profit gene fragment was cut down. The restriction enzyme used the combination of Nde I and Hind III, mixture to 109 shares of competent cel Escherichia coli JM, and it is competent cel Escherichia [0027] [Example 3] Restriction enzyme processing was performed about each pT7 blue plasmid 100 mug ml-1 ampicillin sodium and 100 LB agar medium containing muM IPTG and 0.004 % Xwhich uses the genomic DNA of the strain as mold was performed, and the colony amplified by DNA containing the manifestation system construction PPIase gene of PPIase, and the PPIase shares of competent cel Escherichia coli JM, respectively. It is the suspension of these strain (PERKIN-ELMER). As a result of comparing this array with the base sequence of the genomic respectively. The cut gene fragment is the pET21a plasmid DNA which carried out restriction **** recombination Escherichia coli of PPlase. It is a 2xYT culture medium to the Erlenmeyer enzyme processing beforehand by 2% agarose gel after dissociating and extracting. (Novagen) Gal was inoculated, it cultivated at 37 degrees C, and about the obtained White colony, PCR [0026] Transformation was carried out by adding the above-mentioned ligation liquid to 109 After collecting PT7 plasmid DNA from an electropositive colony, the base sequence of the DNA of pie ROKOKKASU HORIKOSHI on a database, or METANOKOKKASU YANASHII. it checked that it was indifferent from the array of a PPlase gene. The amino acid sequence presumed from it by the array number 1 and the array number 3, respectively in the base ml-1, pH7.5) 700 ml was put in and two to recombination Escherichia coli 3 platinum loop sequence of the PPlase gene of pie ROKOKKASU HORIKOSHI and METANOKOKKASU reaction (a primer being T7 promotor primer and a U-19 reverse primer) using BIG Dye coli about this plasmid. BL21 Transformation was carried out to the stock (DE3). YANASHII is shown in the array number 2 and the array number 4, respectively.

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the Escherichia coli with which a molecular weight marker and a lane 2 discover the fungus body In addition, the lane 1 in drawing 3 makes a sample the Escherichia coli non-refined fungus body discover and case of the Escherichia coli which made (the <u>drawing 3</u> lane 4) and PHFK discover. (10000 rpmx10min). The obtained fungus body was suspended in 25 mM HEPES buffer-solution overnight. Column purification was repeated in order of the anion-exchange chromatography of (110 rpm) at 35 degrees C for 24 hours, fungus bodies were collected in centrifugal separation crushing liquid of wild type Escherichia coli, and the lane 3 made MJFKL discover, and a lane 5 When refined fungus body crushing liquid was separated by SDS-PAGE, (the drawing 3 lane 6) a-d which showed the fungus body to the following after ultrasonic crushing, and gel filtration. crushing liquid which made PHFK discover, and the non-refined fungus body crushing liquid of containing a PHFK gene or a MJFKL gene was inoculated. After carrying out rotation culture (pH6.8) 20 ml containing 1 mM EDTA, and carried out cryopreservation at -20 degrees C was detected as a band also with single case of the Escherichia coli which made MJFKL

liquid 100 % rate of flow : [1] ml min-1b. HiLoad 26/60 Superdex 200pg column () [26 mm x 60 cm; Amersham] Pharmacia Eluate : 100 mM Sodium phosphate buffer solution () [pH7.0;] 0.15 included 25 mM HEPES-KOH The buffer solution (0(pH 6.8)-10 min: B liquid 0 % --) 10-60 min: KOH The buffer solution (pH 6.8) B liquid: 0.5 M NaCl included 25 mM HEPES-KOH The buffer [0029] a. DEAE Toyopearl column A (16 mm x 60 cm; TOSOH Co., Ltd.) liquid: 25 mM HEPES-TOSOH Co., Ltd.) liquid: 25 mM HEPES-KOH The buffer solution (pH 6.8) B liquid: 0.5 M NaCl solution (0(pH 6.8)-300 min. B liquid 0-100% of straight-line gradient ---) 300-420 min. The B M NaCl content rate of flow: 3 ml min-1c. TSKgel SuperQ-5PW column A (7.5 mm x 7.5 cm; SWXLcolumn eluate (7.5 mm x 30 cm; TOSOH Co., Ltd.) : 100 mM Sodium phosphate buffer solution (pH7.0; 0.15 M NaCl content) Rate of flow: 0.5 ml min-1[0030] [Example 5] Protein straight-line gradient rate-of-flow [B liquid 0-100% of]: -- 1 ml min-1d, TSKgel G3000 thermal denaturation depressor effect of PHFK (1)

Escherichia coli which made PHFK discover was carried out, and supernatant liquid was obtained the quantum of the residual protein concentration of supernatant liquid was carried out (drawing electrophoresis which used acrylamide gel 16% about supernatant liquid (drawing 4). Moreover, As contrast, wild type Escherichia coli was used instead of the Escherichia coli which made heat-treated for 30 minutes in the temperature requirement of 20 - 100 **. The protein which according to centrifugal separation. Protein concentration was carried out in 2.5mg/ml, and it The effect of PHFK to the thermal denaturation of the fungus body crushing liquid digestive 5). As contrast, wild type Escherichia coli was used instead or the Escheriona oun minor PHFK discover, and detection and the quantum of the protein in supernatant liquid were liquor of Escherichia coli (BL21 DE3) was considered. That is, ultrasonic crushing of the remains in supernatant liquid was detected after centrifugal separation by the SDS

Escherichia coli did not condense, but it remained in supernatant liquid. On the other hand, when temperature, and protein hardly remained in supernatant liquid at the time of high temperature PPIase did not exist, the amount of detection protein decreased with the rise of processing performed. [0031] As shown in <u>drawing 4</u> and <u>drawing 5</u> , when PPlase existed, the protein of most processing.

 ${\sf Img/ml}$, and PHFK was heat-treated for 30 minutes in the temperature requirement of ${\sf 20-100}$ The protein which remains in supernatant liquid as contrast, without adding PHFK was detected. **. The protein which remains in supematant liquid was detected after centrifugal separation by supernatant liquid was obtained according to centrifugal separation. Protein concentration was carried out in 2.1mg/ml, and further, it added so that it might become the last concentration of remained, but when PPlase was not added, the amount of the protein detected was decreasing the SDS electrophoresis which used acrylamide gel 16% about supernatant liquid (drawing 6). 0033] As shown in drawing 6, when PPlase was added, the protein of most Escherichia coli Ultrasonic crushing of the wild type Escherichia coli (BL21 DE3) was carried out, and [0032] [Example 6] Protein thermal denaturation depressor effect of PHFK (2) with the rise of processing temperature.

http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje

nvention is [playback of denatured protein, stabilization of a protein reagent, and a pan] useful Effect of the Invention] By this invention, the denaturation of the protein by heat, an organic production can be controlled according to this denaturation depressor effect. In addition, this solvent, etc. can be controlled, and irreversible condensation formation can be prevented. Moreover, the inclusion body formation which poses a problem in recombination protein co retrieval of a new immunosuppresant and a physiological active substance.

[Layout Table]

30 aaa gaa gag aga ata tat aat cca aag ggg atc tac ggtcca gtt cca 144 Lys Glu Ala Gly Ile TyrAsn Pro Lys Gly Ile Tyr Gly Pro Val Pro 35 40 45 ata atc gtc gga gct ggt cac gtc att tct gga tta gac aag agg ctg192 Ile Ile Val Gly Ala Gly His Val Ile Ser Gly Leu Asp Lys Arg Leu 50 55 60 gta gga ttt agaaag cag ggg ata gtt cca tto cca gga tta gaa gta 336 Gly Gln Phe Arg Lys Gln Gly Ile Val Pro Phe Pro Gly Leu Glu Val 100 105 110 gaa gtc acg act gacaat gga agg aag atg aaa ggt agg gta att aca 384 Glu ValThr Thr Asp Asn Gly Arg Lys Met Lys Gly Arg Val Ile Thr 115 120 125 gta 2.0<210> 1<211> 774<212> DNA<213> Pyrococcus-horikoshii<220> <221> CDS <222> (1) .. (771) act tac gaa gaa gtg gcc 96 Val Lys Glu Thr Gly Gln lle Phe Asp Thr Thr Tyr Glu Glu Val Ala 2025 ctt gaa gta gga aag aag tac aco tta gag gtt coa coa gag 240 Val Gly Leu Glu Val Gly Lys Lys Tyr Thr Leu Glu Val Pro Pro Glu 65 70 75 80gaagga tttgga cta agg gat coc aag otg att aag gta tto acg atg 288 Glu Gly Phe Gly Leu Arg Asp Pro Lys Leu Ile Lys Val Phe Thr Met 85 90 95 gga caa Asp Lys Arg Leu 50 55 60 Val Gly Leu Glu Val Gly Lys Lys Tyr Thr Leu Glu Val Pro Pro Glu 65 70 75 80 Glu Gly Phe Gly Leu Arg Asp Pro LysLeu Ile Lys Val Phe Thr Met 85 90 95 Gly Gln Phe Arg Lys Gln Gly Ile Val Pro Phe Pro Gly Leu Glu Val 100 105 110 Glu Val Thr Thr AspAsn Gly Arg Lys Met Lys Gly Arg Val Ile Thr 115 120 125 Val Ser Gly Gly Arg Val Arg Val Asp Phe Asn His Pro Leu Ala Gly 130 135 140 Lys Thr Leu Ile Tyr Glu Val Glu Ile Val Glu Lys Ile Glu Asp Pro 145 150 155 160 Ile Glu Lys Ile Lys Ala Leu Ile Glu-Leu-Arg-Leu-Pro-Met-Ile-Asp 165 170 SEQUENCE-LISTING <110> MARINE BIOTECHNOLOGY INSTITUTE CO., LTD.<120> METHOD-FOR-RETARDING DENATURATION-OF-PROTEIN<130> P00-0472<160> 8 <170> Patentin Ver. (400) 1 atg aag gtg gag agg gga gat gtt atc agg ctc cac tat acc ggt agg 48 Met Lys Val Glu Arg Gin Asp Val Asp Pro Lys Thr Leu Ile Leu Gly Giu Ile 195 200 205 Leu Leu Giu Ser Asp Ile Lys Phe Leu Gly Tyr Giu Lys Val Giu Phe 210 215 220 Lys Pro Ser Val Giu Giu Leu Leu Arg Pro Lys cta ttt gat aca act aac gaa gaa ttg gct aaa aaa gag 96 Asp Gly Lys Leu Phe Asp Thr Thr Asn Glu Glu Leu Ala Lys Lys Glu 20 25 30 ggg att tac aac cct goa atg atttat ggt oot gtt got ato ttt gct $\langle 221 \rangle$ CDS $\langle 222 \rangle$ (1) .. (693) $\langle 400 \rangle$ 3 atg gta gaa aag ggt aaa atg gta aag att agc tat gac gga tac 160ata gag aag ata aaa gco cta ata gag otg agg tta oca atg ato gat 528 lle Glu Lys lle Lys Ala Leu lle Glu Leu Arg Leu Pro-Met-lle-Asp 165 170 175 agg gat aag gta ata-atc-gaa-gtt-gag gaa ago gga ggt agg gttaga gtt gat ttt aac cac ccc cta gcc gga 432 Val Ser Gly Gly Arg Val Arg Val Arg Val Asp Phe Asp Phe Asn His Pro Leu Ala Gly 130 135 140 aaa acc ctt atttat gag gtggag att gtt gag aag atc ggatat gag aag gtt gaa ttt 672 Leu Leu GluSer Asp Ile Lys Phe Leu Gly Tyr Glu Lys Val Glu Phe 210 215 220 aaa oot agt gtt gaa gag ttgttg agg ooc aag oag gaa gaa ooc gtt 720 Lys Pro Ser Val aag gat gtt aag gta aac 576 Arg Asp Lys Val Ile Ile-Glu-Val-Gly-Glu Lys Asp Val Lys Val-Asn 180 185 190 ttt ggt gagcaa gat gtt gat cocaag acg ctg atc ctg gga gaa att 624 Phe Gly Glu Gln Asp Val Asp Pro Lys Thr Leu Ile Leu Gly Glu Ile 195 200 205 ctt ttg gag agt gat att aaa ttc ctg 175 Arg Asp Lys Val Ile Ile Glu Val Gly Glu Lys Asp Val Lys-Val-Asn 180 185190 Phe Gly Glu Gin Glu Glu Pro Val 225 230 235 240 Glu Glu Lys Lys Glu Glu Glu Glu Glu Ser Glu Glu Ala Gln Ser 245 250 255Ser (210) 3 (211) 696 (212) DNA (213) Methanococcus jannaschi(220) Gly Asp Val Ile Arg Leu His Tyr Thr Gly Arg 1 5 10 15 gtt aaa gag act gga caa ata ttt gac acc Asn Pro Lys Gly Ile Tyr Gly Pro Val Pro 35 40 45 Ile Ile Val Gly Ala Gly His Vallle Ser Gly Leu gtt 48 Met Val Glu Lys Gly Lys Met Val Lys Ile Ser Tyr Asp Gly Tyr Val 1 5 10 15 gat gga aaa gaa gat cca 480 Lys Thr Leu Ile Tyr Glu Val Glu Ile Val Glu Lys Ile Glu Asp Pro145 150 155

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2007/08/02

Arg Val Leu Val 115 120 125 gat tit aac cat gaa tta gct gga aaa gag gta aaa tat agg ata aaa 432 Asp Phe Asn His Glu Leu Ala Gly Lys Glu Val Lys Tyr Arg Ile Lys 130 135 140 att gaa gaa gtt gtt aaa gttagc ttt gtt gag aca ttt gaa aga aaa aag 672 Glu Asp Ala GluLys Val Ser Phe Val Glu Thr Phe Glu Arg Lys Lys 210 215 220 gaa act aaa gaa gag aac aaataa 696 Glu Thr Lys Glu Glu Asn Lys 225 230 <210> 4 <211> 231<212> PRT <213> Methanoccccus jannaschi <400> 4 Met Val Glu Lys Gly Lys Met Val Glu Lys Gly Lys Met Val Lys Gly Lys Met Val Lys Gly Lys Gly Lys Met Val Lys Gly Lys Gly Lys Leu Phe Asp Thr ThrAsn Glu Glu Leu Ala Lys Lys Glu 20 25 30 Gly Ile Tyr Asn Pro AlaMet Ile Tyr Gly Pro Val Ala lie Phe Ala 35 40 45 Gly Glu Gly Gln Val LeuPro Gly Leu Asp Glu Ala lie Leu Glu Met 50 55 60 Asp Val GlyGlu Glu Arg Glu Val Val Leu Pro Pro Glu Lys Ala Phe 65 70 7580 Gly Lys Arg AspPro Ser Lys Ile Lys Leu Ile Pro Leu Ser Glu Phe 85 90 95 Thr Lys Arg Gly Ile Lys Pro Ile Lys Gly Leu Thr Ile Thr Ile Asp 100 105 110 Gly Ile Pro Gly Lys Ile Val Ser Ile Asn—Ser—Gly—Arg—Val Leu Val 115 120 125 Asp Phe Asn His Glu Leu Ala Gly Lys—Glu—Val—Lys—Tyr—Arg—Ile—Lys 130 135 140 Ile Glu Glu Val Val Asp Asp Lys Lys-Asn Ile Val Lys Glu Ile Val 145 150 155160 Lys Met Tyr Val Pro Arg Leu Ser Asp Val Lys Val Thr Ile Arg Asn 165 170 175 Gly Thr Val Lys Ile Glu Leu Pro Glu Phe Ala Pro Phe Ile Pro Asn 180 185 190 Ile Gln Thr Ala Lys Met Ala Ile Ala Asn Leu Ser Glu Phe 85 90 95 aca aaa aga ggaatt aag coa ata aaa gga tta acc ata act att gat 336 Thr Lys Arg Gly lle Lys Pro-lle-Lys-Gly-Leu-Thr-lle-Thr-lle-Asp 100 105 110 gga att cot gga aaa att gtt agc ata aac agt gga aga gtt tta gtc 384 Gly lle Pro Gly Lys lle Val Ser lle Asn Ser Gly ata gaa ttg oot gaa ttt got ooa ttt att ooa aac 576 Gly Thr Val Lys Ile Glu Leu Pro Glu Phe Ala Pro Phe Ile Pro Asn 180 185 190 att oaa aca got aagatg got att got aac gaa ata ttg aag aga tta 624 Ile Gln Thr Ala Lys Met Ala Ile Ala Asn Glu Ile Leu Lys Arg Leu 195 200 205 gaa gat got gaa 144 Gly lle Tyr Asn Pro Ala Met Ile Tyr Gly Pro Val Ala Ile Phe Ala 35 40 45 gga gaa gga caa gta ttacct gga tta gac gaa gcc ata tta gaa atg 192 Gly Glu Gly Gln Val Leu Pro Gly Leu Asp Glu Ala gat gat aaa aag aat att gta aaa gaa att gta 480 lle Glu Glu Val Val Asp Asp Lys Lys Asn Ile Val Lys Glu Ile Val145 150 155 160aaa atg tat gtt cca aga ttgagtgat gta aaa gta act atc aga aat 528 Lys Met Tyr Val Pro Arg Leu Ser Asp Val Lys Val Thr Ile Arg Asn 165 170 175gga aca gtt aag toa aag ata aaa tta ato coa tta toa gaa ttt 288 Gly Lys Arg Asp Pro Ser Lys Ile Lys Leu Ile Pro Glu Ile Leu Lys Arg Leu 195 200 205 Glu Asp Ala Glu Lys Val Ser Phe Val Glu Thr Phe Glu Arg Lys Lys 210 215 220 Glu Thr Lys Glu Glu Asn Lys 225 230 <210> 5 <211> 30<212> DNA<213> lle Leu Glu Met 50 55 60 gat git ggt gaggaa aga gaagit git tta cot coa gag aaa got tit 240 Asp Val Gly Glu Glu Arg Glu Val Val Leu Pro Pro Glu Lys Ala Phe 65 70 75 80ggt aag aga gac coa 28<212> DNA<213> Artificial Sequence <400> 7ccatatggta gaaaagggta aaatggta 28 <210> 8 (211) 29(212) DNA (213) Artificial Sequence (400) 8 ggggatcott attigitoto ttotttagt 29 DNA<213> Artificial Sequence <400> 6ggaagctttt aagaggattg cgcctcttc 29<210> 7 <211> Artificial Sequence (400) 5cccatatgaa ggtggagagg ggagatgtt 30(210) 6 (211) 29(212)

[Translation done.]

* NOTICES *

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1. This document has been translated by computer. So the translation may not reflect the original

2.*** shows the word which can not be translated.

3.In the drawings, any words are not translated.

DESCRIPTION OF DRAWINGS

[Brief Description of the Drawings]

[Drawing 1] Drawing showing the amino acid sequence presumed from the base sequence of a PHFK gene, and it.

[Drawing 2] Drawing showing the amino acid sequence presumed from the base sequence of a MJFKL gene, and it.

[Drawing 3] Electropherogram of the fungus body crushing liquid of the Escherichia coli which discovers MJFKL or PHFK

[Drawing 4] Electropherogram of the fungus body crushing liquid supernatant liquid under PHFK

existence and nonexistence

<u>[Drawing 5]</u> Drawing showing the amount of protein in the fungus body crushing liquid
supernatant liquid under PHFK existence and nonexistence

<u>[Drawing 6]</u> Electropherogram of the fungus body crushing liquid supernatant liquid at the time of
PHFK addition and additive-free

[Translation done.]